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Our Docket No. CMCC 779

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### MESSAGE:

## IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant:

Samy Ashkar

Serial No.:

09/981,845

Art Unit:

1647

Filed:

October 18, 2001

Examiner:

Regina M. Deberry

For:

OSTEOPONTIN-COATED SURFACES AND METHODS OF USE

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## IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Appellants:

Samy Ashkar and Jairo Salcedo

Serial No.:

09/981,845

Art Unit:

1647

Filed:

October 18, 2001

Examiner:

Regina M. Deberry

For:

OSTEOPONTIN-COATED SURFACES AND METHODS OF USE

Mail Stop Appeal Brief-Patents Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

## APPEAL BRIEF

Sir:

This is an appeal from the final rejection of claims I-6 in the Office Action mailed February 13, 2004, in the above-identified patent application. A Notice of Appeal was mailed on June 14, 2004 (there is an error in the Advisory Action mailed June 28, 2004). The Commissioner is hereby authorized to charge \$165.00, the fee for the filing of this Appeal Brief for a small entity, to Deposit Account No. 50-3129. It is believed that no additional fee is required with this submission. However, should an additional fee be required, the Commissioner is hereby authorized to charge the fee to Deposit Account No. 50-3129.

# (1) REAL PARTY IN INTEREST

45049567v1

CMCC 779 078856/00047 U.S.S.N. 09/981,845 October 18, 2001 Filed: APPEAL BRIEF

The real party in interest of this application is Children's Medical Center Corporation in Boston, MA, the assignce of record; and the licensee of record OraPharma, Inc. in Warminster, PA.

#### RELATED APPEALS AND INTERFERENCES (2)

There are no related appeals or interferences known to appellant, the undersigned, or appellant's assignce which directly affects, which would be directly affected by, or which would have a bearing on the Board's decision in this appeal.

#### STATUS OF CLAIMS ON APPEAL **(3)**

Claims 1-6 are pending. Claims 1-6 are on appeal. Claims 7-18 were cancelled in an Amendment filed on November 21, 2003. The text of each claim on appeal, as pending, is set forth in an Appendix to this Appeal Brief.

#### STATUS OF AMENDMENTS **(4)**

An amendment after final rejection was mailed on May 11, 2004. In the Advisory Action mailed June 28, 2004, the Examiner indicated that this amendment would be entered. An appendix sets forth the claims on appeal.

#### SUMMARY OF THE INVENTION **(5)**

The claims are drawn to isolated active osteopontin fragments and osteopontin-derived peptide fragments that have cell-attachment and cell-spread activity (page 7, line 23 to page 8, line 12). The peptide fragments may be used to increase cell attachment to a material, as well as enhance cell spread on the material (page 11, lines 9-18). The material is suitable for use on a material which is implanted into a patient to enhance cell-attachment and cell-spread activity and 45049567v1

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APPEAL BRIEF

thereby integration of the implant, for example, for use in treatment of periodontal disease (page 10, lines 16-23). Claim 1 is directed to an osteopontin-derived peptide fragment comprising an amino acid sequence selected from the group consisting of SEQ 1D NO:7, SEQ 1D NO:8, SEQ 1D NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, and SEQ ID NO:15 (page 8, lines 7-26 and page 12, lines 4-13). Claim 2 is directed to the peptide fragment of claim 1, wherein the peptide increases cell attachment to a material and increases cell spread (page 8, lines 11-12 and page 53, lines 12-17). Claim 3 is directed to the peptide fragment of claim 2, wherein the peptide binds to at least one receptor on a cell surface. Claim 4 is directed to the peptide fragment of claim 3, wherein the receptor(s) is an integrin. Claim 5 is directed to the peptide fragment of claim 4, wherein the integrin(s) is  $\alpha_v \beta_3$ ,  $\alpha_v \beta_5$ ,  $4\beta_1$ ,  $2\beta_1$ . VCAM, ICAM CD44, or  $V_3 V_x$ . Support for claims 3, 4, and 5 can be found on page 3, line 27 to page 4, line 14 and page 53, lines 17-21. Claim 6 is directed to the peptide fragment of claim 3 wherein the cell is an osteoprogenitor cell, tumor cell, macrophage, periosteal cell, endothclial cell, epithelial cell, cosinophil, stem cell, limited potential precursor cell, precursor cells committed precursor cell, or differentiated cell (page 8, line 29 to page 9, line 2).

# (6) ISSUES ON APPEAL

The issues presented on appeal are:

(1) whether claims 1-6 are enabled under 35 U.S.C. § 112, first paragraph.

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#### ARGUMENTS **(7)**

# (a) The Claimed Invention

The claims are directed to active osteopontin-derived peptide fragments and their use in and/or on materials to increase cell attachment and cell spread activity. The peptides may be used to coat, for example, a surgical implant where cell attachment and growth on the implant are desirable. The peptide fragments comprise the sequences VFTPVVPTVDTYDGRGDSVVYGLRSKSKKFRRP (SEQ ID NO. 7), RSRRATEVFTPVVPTVDTYDGRGDSVVYGLRSKSKKFRRP (SEQ ID NO:8), SDELVTDFPTDLPATEVFTPVVPTVDTYDGRGDSVVYGLRSKSKKFRRP (SEQ ID NO:9), RSRRATEVFTPVVPTVDTYDGRGDSVVYGRRSKSKKFRRP (SEQ ID NO:10), RSRRATEVETPVVP1 VDTYDGRGDSVVYGRRSKSKKFRRPAGAAGGPAGPAG PAGPAGPAGPA (SEQ ID NO:11), RSRRVFTPFIPTESANDGRGDSVAYGLKSKSKKFRR (SEQ ID NO:12), DTFTPIVPTVDVPNGRFDSLAYGLKSKSKKFQ (SEQ ID NO:13), RSRRATEVFTPVVPTVDTYDGRADSVVYGRRSKSKKFRRP (SEQ ID NO:14), and acetyl-RSRRATEVFTPVVPTVDTYDGRGDSVVYGLRSKSKKFRRP (SEQ ID NO:15).

The osteopontin-derived peptide fragments increase cell binding and spread by binding to integrins, such as  $\alpha_V \beta_3$ ,  $\alpha_V \beta_5$ ,  $4\beta_1$ ,  $2\beta_1$ , VCAM, ICAM CD44,  $V_3 V_x$ , on the surface of cells. The peptide fragments may be used to modulate a number of different cell types, including osteoprogenitor cells, tumor cells, macrophages, periosteal cells, endothelial cells, epithelial cells, eosinophils, stem cells, limited potential precursor cells, precursor cells, committed precursor cells, and differentiated cells.

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4

CMCC 779 078856/00047 The Legal Standard

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The peptides have numerous applications, but principally in tissue repair or regeneration, for example, when coated onto a titanium material and used in the treatment of periodontal disease to enhance bone regrowth.

# (b) Rejection of claims 1-6 Under 35 U.S.C. § 112, first paragraph

The Court of Appeals for the Federal Circuit (CAFC) has described the legal standard for enablement under § 112, first paragraph, as whether one skilled in the art could make and use the claimed invention from the disclosures in the patent coupled with information known in the art as of the date of filing, without undue experimentation (See, e.g., Amgen v. Hoechst Marion Roussell 314 F.3d 1313 (Fed. Cir. 2003; Genentech, Inc. v. Novo Nordisk A/S, 108 F3d at 165, 42 USPQ2d at 1004 (Fed. Cir. 1997) (quoting In re Wright, 999 F.2d 1557, 1561, 27 USPQ2d 1510, 1513 (Fed. Cir. 1993); See also In re Fisher, 427 F.2d at 839, 166 USPQ at 24; United States v. Telectronics, Inc., 857 F.2d 778 (Fed. Cir. 1988); In re Stephens, 529 F.2d 1343 (CCPA 1976)). The fact that experimentation may be complex does not necessarily make it undue, if the art typically engages in such experimentation (M.I.T. v. A.B. Fortia, 774 F.2d 1104 (Fed. Cir. 1985)). As affirmed by the Court in Spectra-Physics, Inc. v. Coherent, Inc., 827 F.2d 1524 (Fed. Cir. 1987), a patent need not teach, and preferably omits, what is well known in the art.

Whether the disclosure is enabling is a legal conclusion based upon several underlying factual inquiries. See In re Wands, 858 F.2d 731, 735, 736-737, 8 USPQ2d 1400, 1402, 1404 (Fed. Cir. 1988). As set forth in Wands, the factors to be considered in determining whether a claimed invention is enabled throughout its scope without undue experimentation include the **CMCC 779** 45049567v1 5

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U.S.S.N. 09/981,845 Filed: October 18, 2001

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quantity of experimentation necessary, the amount of direction or guidance presented, the presence or absence of working examples, the nature of the invention, the state of the prior art, the relative skill of those in the art, the predictability or unpredictability of the art, and the breadth of the claims. In cases that involve unpredictable factors, "the scope of the enablement obviously varies inversely with the degree of unpredictability of the factors involved." *In re Fisher*, 427 F.2d 833, 839, 166 USPQ 18, 24 (CCPA 1970). The fact that some experimentation is necessary does not preclude enablement; what is required is that the amount of experimentation "must not be unduly extensive." *Atlas Powder Co.*, v. E.I. DuPont De Nemours & Co., 750 F.2d 1569, 1576, 224 USPQ 409, 413 (Fed. Cir.1984). There is no requirement for examples.

## Analysis

A proper analysis of the *Wands* factors shows that claims 1-6 satisfy the enablement requirement. The quantity of experimentation necessary to make and use the claimed peptides is **not undue.** The claims are directed to ostepontin-derived peptide fragements comprising SEQ ID NO:7, SEQ ID NO:3, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, or SEQ ID NO:15. These sequences are well known. The amino acid sequence and structure of osteopontin, from which the peptide fragments are derived, are well known. One skilled in the art would have no difficulty making short peptides synthetically, or longer peptides using a portion of the nucleotide sequence encoding osteopontin. The point of novelty is the identification of the amino acid sequence in a very large protein which has the desired activity, and that this activity is retained even in a very small peptide relative to the huge

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protein from which it is derived. The specification describes how to coat the peptides to a material (page 13, line 14 to page 14, line 21) and describes the types of materials that may be coated (page 10, lines 16-23 and page 14, lines 22-28). The specification describes the cell types that may be regulated using the osteopontin-derived peptides fragments (page 8, line 29 to page 9, line 2) and that the peptides bind integrin receptors on the surface of these cells (page 3, line 27 to page 4, line 14).

Although there is no requirement for examples, Example 12 and Table 8 on pages 53-55 of the originally filed application, demonstrate that each of SEQ ID NO:15, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, or SEQ ID NO:14 binds to osteoprogenitor cells and significantly increases cellular attachment and spread over the control. In addition, Example 12 and Table 8 illustrate that antibodies to integrins (i.e.,  $\alpha_v \beta_3$ ) inhibit the percentage of attached cells and cell spread induced by the peptides (i.e., SEQ ID NO: 15), indicating that the peptides interact with integrins.

The guidance in the specification and ease in carrying out the assays, as shown in the PAGE 10/10 \* RCVD AT 8/16/2004 10:25:09 PM [Eastern Daylight Time] \* SVR:USPTO-EFXRF-1/0 \* DNIS:8729306 \* CSID:706 283 7737 \* DURATION (mm-ss):03-42